



Ortho-Clinical Diagnostics

a Johnson & Johnson company

100 Indigo Creek Drive
Rochester, New York 14626-5101

510(k) Summary

This summary of 510(k) safety and effectiveness information is being submitted in accordance with the requirements of SMDA 1990 and 21 CFR 807.92.

The assigned 510(k) number is: K062094

**1. Submitter
name,
address,
contact**

Ortho-Clinical Diagnostics, Inc.
100 Indigo Creek Drive
Rochester, New York 14626-5101
(585) 453-4253
Contact Person: Darlene Phillips

**2. Preparation
Date**

September 28, 2006

3. Device name

Trade or Proprietary Names:

VITROS Chemistry Products PCP Reagent
VITROS Chemistry Products Calibrator Kit 26
VITROS Chemistry Products FS Calibrator 1
VITROS Chemistry Products DAT Performance Verifiers I, II, & V

Common Names:

Phencyclidine (PCP) assay and controls

Classification Names:

Phencyclidine test system (862.3100) Class II
Clinical toxicology calibrators (862.3200) Class II
Clinical toxicology control material (862.3280) Class I, VITROS DAT
Performance Verifiers are assayed controls, so they meet the
reserved criteria under Section 510(l) of the Food, Drug and
Cosmetic Act.

**4. Predicate
Devices**

The VITROS Chemistry Products PCP assay is substantially equivalent to the DADE BEHRING Syva[®] EMIT[®] II Plus Phencyclidine Assay.

The VITROS Chemistry Products DAT Performance Verifiers are substantially equivalent to the BIO-RAD Liquichek[™] Urine Toxicology Controls.

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5. Device description

The VITROS PCP assay is a homogeneous enzyme immunoassay that is performed using the VITROS Chemistry Products PCP Reagent with the VITROS Chemistry Products Calibrator Kit 26, VITROS Chemistry Products FS Calibrator 1 and VITROS Chemistry Products FS Diluent Pack 4 (DAT Diluent / DAT Diluent 2) on VITROS 5,1 FS Chemistry Systems.

The VITROS PCP Reagent is a dual chambered package containing ready-to-use liquid reagents that are used to detect phencyclidine in urine. Samples, calibrators, and controls are automatically treated with surfactant (DAT Diluent 2) prior to addition of reagents. Treated sample is added to Reagent 1 containing antibody reactive to phencyclidine, glucose-6-phosphate and nicotinamide adenine dinucleotide (NAD⁺), followed by Reagent 2 containing phencyclidine labeled with the enzyme glucose-6-phosphate dehydrogenase (G6P-DH). The assay is based on competition between phencyclidine in the treated urine sample and phencyclidine labeled with the enzyme glucose-6-phosphate dehydrogenase (G6P-DH) for antibody binding sites. Enzyme activity decreases upon binding to the antibody, therefore the concentration of phencyclidine in the urine sample is directly proportional to measured enzyme activity. Active enzyme converts oxidized nicotinamide adenine dinucleotide (NAD⁺) to NADH, resulting in an absorbance change that is measured spectrophotometrically at 340 nm.

VITROS Chemistry Products Calibrator Kit 26 is prepared from human urine to which drugs of abuse, metabolites of drugs of abuse, organic salt, surfactants, and preservative have been added. VITROS FS Calibrator 1 is prepared from sodium chloride and processed water. VITROS Calibrator Kit 26 is used in conjunction with VITROS FS Calibrator 1 to calibrate VITROS 5,1 FS Chemistry Systems for the qualitative or semi-quantitative measurement of phencyclidine (PCP).

VITROS Chemistry Products DAT Performance Verifiers I, II, and V are prepared from a human urine pool to which analytes, surfactant, and preservative have been added. These are assayed controls used to monitor performance of the VITROS PCP assay on VITROS 5,1 FS Chemistry Systems.

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- 4. Device description**
(continued)
- The VITROS Chemistry Products FS Diluent Pack 4 (DAT Diluent/ DAT Diluent 2) is a common reagent that is used with several drugs of abuse assays to dilute calibrators and samples on the VITROS 5,1 FS System. This is a dual chambered package containing two ready-to-use liquid diluents. DAT Diluent is prepared from human urine to which organic salt, surfactants, and preservative have been added. DAT Diluent 2 is prepared from processed water to which surfactant and preservative have been added.
- The VITROS 5,1 FS Chemistry System is a clinical chemistry instrument that provides automated use of the VITROS Chemistry Products MicroTip[®] and MicroSlides[®] range of products. The VITROS 5,1 FS System was cleared for market by 510(k) premarket notification (K031924).
- 6. Device intended uses**
- VITROS Chemistry Products PCP Reagent:** For *in vitro* diagnostic use only. VITROS Chemistry Products PCP Reagent is used on VITROS 5,1 FS Chemistry Systems for the semi-quantitative or qualitative determination of phencyclidine (PCP) in human urine using a cutoff of 25 ng/mL. Measurements obtained with the VITROS PCP method are used in the diagnosis and treatment of phencyclidine use or overdose.
- The VITROS Chemistry Products PCP assay is intended for use by professional laboratory personnel. It provides only a preliminary test result. A more specific alternative chemical method must be used to confirm a result obtained with this assay. Gas chromatography/mass spectrometry (GC/MS) is the preferred confirmatory method. Clinical consideration and professional judgment should be applied to any drug-of-abuse test result, particularly when evaluating a preliminary positive result.
- VITROS Chemistry Products Calibrator Kit 26:** For *in vitro* diagnostic use only. VITROS Chemistry Products Calibrator Kit 26 is used to calibrate VITROS 5,1 FS Chemistry Systems for the qualitative or semi-quantitative measurement of drugs of abuse.
- VITROS Chemistry Products FS Calibrator 1:** For *in vitro* diagnostic use only. VITROS Chemistry Products FS Calibrator 1 is used in conjunction with VITROS Chemistry Products Calibrator Kits to calibrate VITROS 5,1 FS Chemistry Systems.
- VITROS Chemistry Products DAT Performance Verifiers I, II, and V:** For *in vitro* diagnostic use only. VITROS Chemistry Products DAT Performance Verifiers are assayed controls used to monitor performance of urine drugs of abuse screening assays on VITROS 5,1 FS Chemistry Systems.

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- 7. Comparison to predicate devices** The VITROS Chemistry Products PCP assay and VITROS Chemistry Products DAT Performance Verifiers are substantially equivalent to the DADE BEHRING Syva[®] EMIT[®] II Plus Phencyclidine Assay (K993983) and BIO-RAD Liquichek[™] Urine Toxicology Controls (K022707) (predicate devices) which were cleared by the FDA for in vitro diagnostic use.

The performance of the VITROS PCP assay on the VITROS 5,1 FS Chemistry System was compared to the Syva[®] EMIT[®] II Plus Phencyclidine assay on the Syva[®] 30R Biochemical System. The results demonstrated good agreement between the two immunoassay methods.

The VITROS PCP assay and the VITROS DAT Performance Verifiers have the following similarities to the predicate devices: the same intended use, the same cutoff value of 25 ng/mL, consist of liquid, ready to use reagents, have similar performance characteristics, are used on an automated clinical chemistry analyzer and calibrated against the same drug, phencyclidine.

Table 1 Similarities and differences of the assays performed using the VITROS PCP assay and the VITROS DAT Performance Verifiers and the EMIT[®] Phencyclidine assay and BIO-RAD[®] Liquichek[™] Urine Toxicology Controls.

Device Similarities	
Device Characteristic	Description
Indications for Use	The assays are intended for use in the qualitative and semi-quantitative analysis of phencyclidine in human urine.
Calibration traceability	Phencyclidine with confirmation by GC/MS
Cut-Off value	25 ng/mL
Sample Type	Human Urine
Reagent Format	Liquid ready to use
Antibody source	Sheep polyclonal antibodies reactive to phencyclidine
Calibration traceability	Phencyclidine with confirmation by GC/MS
Calibrator matrix	Human urine
Control matrix	Human urine

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Differences		
Device Characteristic	VITROS PCP assay (New device)	EMIT[®] Phencyclidine assay and Liquichek[™] Urine Toxicology Controls (Predicate devices)
Reportable Range	6.0 – 72.0 ng/mL	8 – 90 ng/mL
Calibrator: Number of levels	Five	Qualitative: Three Semi-quantitative: Five
Calibrator format	Frozen Liquid ready to use	Refrigerated liquid ready to use
Instrumentation	To be used on VITROS 5,1 FS Chemistry Systems	Multiple automated clinical chemistry systems
Control claimed analytes	Cocaine metabolites (benzoylecgonine), benzodiazepines (lorazepam), methadone, amphetamines (d-methamphetamine), opiates (morphine), cannabinoids (11-nor- Δ^9 -THC-9-COOH), phencyclidine and barbiturates (secobarbital).	Methamphetamine, secobarbital, lorazepam, tetrahydrocannabinol (THC), benzoylecgonine, ethanol, lysergic acid diethylamide (LSD), methadone, methaqualone, morphine, (Free), phencyclidine, propoxyphene, nortriptyline and addition of creatinine, pH, specific gravity.
Control: Number of levels	Three	Two

8. Conclusions The data presented in the premarket notification provide a reasonable assurance that the VITROS Chemistry Products PCP Reagent, VITROS Chemistry Products Calibrator Kit 26, VITROS Chemistry Products FS Calibrator 1 and VITROS Chemistry Products DAT Performance Verifiers are safe and effective for the stated intended uses and are substantially equivalent to the cleared predicate devices. Equivalence to the predicates was demonstrated using a commercially available assay along with patient samples.



DEPARTMENT OF HEALTH & HUMAN SERVICES

Public Health Service

Food and Drug Administration
2098 Gaither Road
Rockville MD 20850

Ms. Darlene Phillips
Ortho-Clinical Diagnostics, Inc.
100 Indigo Creek Drive
Rochester, New York 14626-5101

NOV 22 2006

Rc: k062094
Trade/Device Name: VITROS Chemistry Products PCP Reagent
Regulation Number: Unclassified
Regulation Name: Enzyme immunoassay, phencyclidine
Regulatory Class: 510(k) required
Product Code: LCM, DKB, DIF
Dated: September 28, 2006
Received: September 29, 2006

Dear Ms. Phillips:

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments, or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (Act) that do not require approval of a premarket approval application (PMA). You may, therefore, market the device, subject to the general controls provisions of the Act. The general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration.

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to such additional controls. Existing major regulations affecting your device can be found in Title 21, Code of Federal Regulations (CFR), Parts 800 to 895. In addition, FDA may publish further announcements concerning your device in the Federal Register.

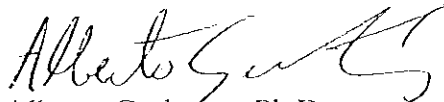
Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal statutes and regulations administered by other Federal agencies. You must comply with all the Act's requirements, including, but not limited to: registration and listing (21 CFR Part 807); labeling (21 CFR Parts 801 and 809); and good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820).

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This letter will allow you to begin marketing your device as described in your Section 510(k) premarket notification. The FDA finding of substantial equivalence of your device to a legally marketed predicate device results in a classification for your device and thus, permits your device to proceed to the market.

If you desire specific information about the application of labeling requirements to your device, or questions on the promotion and advertising of your device, please contact the Office of In Vitro Diagnostic Device Evaluation and Safety at (240) 276-0484. Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21CFR Part 807.97). You may obtain other general information on your responsibilities under the Act from the Division of Small Manufacturers, International and Consumer Assistance at its toll-free number (800) 638-2041 or (301) 443-6597 or at its Internet address <http://www.fda.gov/cdrh/industry/support/index.html>.

Sincerely yours,

A handwritten signature in black ink, appearing to read "Alberto Gutierrez", with a stylized flourish at the end.

Alberto Gutierrez, Ph.D.

Director

Division of Chemistry and Toxicology

Office of In Vitro Diagnostic Device

Evaluation and Safety

Center for Devices and

Radiological Health

Enclosure

Indications for Use

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510(k)
Number (if
known):

K062094

Device Name: VITROS Chemistry Products PCP Reagent

**Indications
for Use:**

For *in vitro* diagnostic use only. VITROS Chemistry Products PCP Reagent is used on VITROS 5,1 FS Chemistry Systems for the semi-quantitative or qualitative determination of phencyclidine (PCP) in human urine using a cutoff of 25 ng/mL. Measurements obtained with the VITROS PCP method are used in the diagnosis and treatment of phencyclidine use or overdose.

The VITROS Chemistry Products PCP assay is intended for use by professional laboratory personnel. It provides only a preliminary test result. A more specific alternative chemical method must be used to confirm a result obtained with this assay. Gas chromatography/mass spectrometry (GC/MS) is the preferred confirmatory method. Clinical consideration and professional judgment should be applied to any drug-of-abuse test result, particularly when evaluating a preliminary positive result.

Prescription Use X
(Part 21 CFR 801 Subpart D)

AND/OR

Over-The-Counter Use
(21 CFR 807 Subpart C)

(PLEASE DO NOT WRITE BELOW THIS LINE-CONTINUE ON ANOTHER PAGE IF NEEDED)

Concurrence of CDRH, Office of In Vitro Diagnostic Devices (OIVD)


Division Sign-Off

Office of In Vitro Diagnostic Device
Evaluation and Safety

510(k) K062094

Indications for Use

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510(k)
Number (if
known):

K062094

Device Name: VITROS Chemistry Products Calibrator Kit 26
VITROS Chemistry Products FS Calibrator 1
VITROS Chemistry Products DAT Performance Verifiers I, II, and V

Indications for Use: For *in vitro* diagnostic use only. VITROS Chemistry Products Calibrator Kit 26 is used to calibrate VITROS 5,1 FS Chemistry Systems for the qualitative or semi-quantitative measurement of drugs of abuse.

For *in vitro* diagnostic use only. VITROS Chemistry Products FS Calibrator 1 is used in conjunction with VITROS Chemistry Products Calibrator Kits to calibrate VITROS 5,1 FS Chemistry Systems.

For *in vitro* diagnostic use only. VITROS Chemistry Products DAT Performance Verifiers are assayed controls used to monitor performance of urine drugs of abuse screening assays on VITROS 5,1 FS Chemistry Systems.

Prescription Use X
(Part 21 CFR 801 Subpart D)

AND/OR

Over-The-Counter Use _____
(21 CFR 807 Subpart C)

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Concurrence of CDRH, Office of In Vitro Diagnostic Devices (OIVD)


Division Sign-Off

Office of In Vitro Diagnostic Device
Evaluation and Safety

(100) K062094